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Please find below and/or attached an Office communication concerning this application or proceeding.

The time period for reply, if any, is set in the attached communication.

		Application No.	Applicant(s)	
Office Action Summary		10/735,959	DRECHSEL ET AL.	
		Examiner	Art Unit	
		MINA HAGHIGHATIAN	1616	
The MAILING DA Period for Reply	ATE of this communication ap	pears on the cover sheet with th	e correspondence address	
WHICHEVER IS LONG  - Extensions of time may be avarafter SIX (6) MONTHS from the  - If NO period for reply is specification.  - Failure to reply within the set of	SER, FROM THE MAILING I ailable under the provisions of 37 CFR 1 e mailing date of this communication. ed above, the maximum statutory period or extended period for reply will, by statu- ce later than three months after the maili	LY IS SET TO EXPIRE 3 MONT DATE OF THIS COMMUNICATI .136(a). In no event, however, may a reply be will apply and will expire SIX (6) MONTHS from the cause the application to become ABANDO and date of this communication, even if timely the	ON.  timely filed  om the mailing date of this communication.  NED (35 U.S.C. § 133).	
Status				
2a)⊠ This action is <b>FIN</b> 3)□ Since this applica	ation is in condition for allowa	April 2008. is action is non-final. ance except for formal matters,   Ex parte Quayle, 1935 C.D. 11,		
Disposition of Claims				
4a) Of the above 5) ☐ Claim(s) is 6) ☑ Claim(s) <u>1-14,16</u> 7) ☐ Claim(s) is 8) ☐ Claim(s) a	claim(s) is/are withdra s/are allowed. 18-20,22-31,38-66,68 and 7	<u>70-95</u> is/are rejected.	cation.	
Application Papers				
10) The drawing(s) file  Applicant may not  Replacement draw	request that any objection to the ing sheet(s) including the corre	er.  cepted or b) objected to by the drawing(s) be held in abeyance. Setion is required if the drawing(s) is examiner. Note the attached Offi	See 37 CFR 1.85(a). objected to. See 37 CFR 1.121(d).	
Priority under 35 U.S.C. §	119			
<ul> <li>12) Acknowledgment is made of a claim for foreign priority under 35 U.S.C. § 119(a)-(d) or (f).</li> <li>a) All b) Some * c) None of:</li> <li>1. Certified copies of the priority documents have been received.</li> <li>2. Certified copies of the priority documents have been received in Application No</li> <li>3. Copies of the certified copies of the priority documents have been received in this National Stage application from the International Bureau (PCT Rule 17.2(a)).</li> <li>* See the attached detailed Office action for a list of the certified copies not received.</li> </ul>				
Attachment(s)  1) Notice of References Cited 2) Notice of Draftsperson's Pa 3) Information Disclosure State Paper No(s)/Mail Date 04/2	ttent Drawing Review (PTO-948) ement(s) (PTO/SB/08)	4) Interview Summa Paper No(s)/Mail 5) Notice of Informa 6) Other:		

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## **DETAILED ACTION**

Receipt is acknowledged of the Amendments, Response and an IDS filed on 04/21/08. Claims 1, 18, 19, 53 and 70 have been amended, claims 15, 17, 67 and 69 have been cancelled and no new claims have been added. Accordingly, claims 1-14, 16, 18-20, 22-31, 38-66, 68 and 70-95 remain pending.

## Claim Rejections - 35 USC § 103

The following is a quotation of 35 U.S.C. 103(a) which forms the basis for all obviousness rejections set forth in this Office action:

(a) A patent may not be obtained though the invention is not identically disclosed or described as set forth in section 102 of this title, if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains. Patentability shall not be negatived by the manner in which the invention was made.

The factual inquiries set forth in *Graham* v. *John Deere Co.*, 383 U.S. 1, 148 USPQ 459 (1966), that are applied for establishing a background for determining obviousness under 35 U.S.C. 103(a) are summarized as follows:

- 1. Determining the scope and contents of the prior art.
- 2. Ascertaining the differences between the prior art and the claims at issue.
- 3. Resolving the level of ordinary skill in the pertinent art.
- 4. Considering objective evidence present in the application indicating obviousness or nonobviousness.

Claims 1-14, 16, 18-20, 22-31, 38-66, 68 and 70-93 are rejected under 35 U.S.C. 103(a) as being unpatentable over Freund et al (DE 19653969 as evidenced by US 2001/0008632).

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Freund teach pharmaceutical preparations in the form of aqueous solutions for the production of propellant-free aerosols for inhalation for the therapy of obstructive lung diseases. Pharmaceuticals intended for inhalation are dissolved in an aqueous or ethanolic solution or a solvent mixture of ethanol and water. The amount of dissolved pharmaceutical in the preparation is between 0.001 and 30%, and preferably between 0.05 and 3%. All substances which are suitable for application by inhalation and which are soluble in the specified solvent can be used as pharmaceuticals in the new preparation. Of especial interest are betamimetics, anticholinergics, antiallergic, antihistamines and steroids, as well as combinations of these active ingredients (sections [0001] to [0007]).

Freund teaches that addition of an effective amount of a complexing agent, such as, **EDTA**, **citric acid**, **ascorbic acid and their salts**, and more especially disodium salt of ethylenediaminetetraacetic acid, eradicates the problem of spray anomalies. The effective quantity of complexing agent Na-EDTA is between 10 and 100 mg/100 ml. Also if necessary, ethanol can be added to increase solubility up to 70% by volume. Other adjuvants such as preservatives, especially benzalkonium chloride can be added in amounts of between 8 and 12 mg/100 ml (sections [0009] to [0013]).

Freund discloses a list of compounds which can be used as active ingredients, singly or in combination, in the aqueous pharmaceutical preparation. In individual cases, it may be required to add a higher quantity of ethanol or a solution mediator to improve solubility. The list includes; **tiotropium bromide**, budesonide, beclomethasone,

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disodium cromoglycate, etc. The solutions are set to a pH of 3.2 to 3.4 with 0.1 or 1 N **HCL** in 100 ml of finished preparation (see sections [0014] to [0046] and [0055]).

It would have been obvious to one of ordinary skill in the art at the time the invention was made to have modified the formulations comprising tiotropium, solvent, an acid and other additives such as benzalkonium chloride with a pH level of 3.2 to 3.4 as taught by Freund et al to prepare the same formulations with a pH of less than 3.2 or 2.0 to 3.0 with a reasonable expectation of successfully preparing safe and stable formulations. In another word, the claims would have been obvious because the technique for improving a particular product was part of the ordinary capabilities of a person of ordinary skill in the art, in view of the teaching of the technique for improvement in other situations. In this situation the improvement is lowering pH levels. One of ordinary skill is well aware that by adjusting the concentration of the acid the pH levels would be adjusted. Freund et al teach that low pH levels are suitable for the said formulations, and one could further lower the pH levels to test for stability.

Claims 1-14, 16, 18-20, 22-31, 38-66, 68 and 70-93 are rejected under 35 U.S.C. 103(a) as being unpatentable over Jager et al (WO 9413262) in view of Bozung et al (DE 19921693 as evidenced by US patent 6,433,027).

Jager et al teach stabilized medicinal aerosol solution formulations comprising

medicaments that degrade or decompose by interaction with solvents or water. The most preferred examples of the medicaments for use in the aerosol solution formulations include ipratropium bromide, tiotropium bromide and fenoterol hydrobromide (see page 8, lines 3-9). Suitable solvents include ethanol and water (see pages 9-10 and examples). One or more acids are added to effect the rate of degradation of the medicament and adjust the pH. Such acids include inorganic acids such as hydrochloric acid and nitric acid or organic acids such as ascorbic acid and citric acid. In aqueous solution the rate of hydrolysis and esterification is typically pH dependent. In aqueous solution, the degradation of ipratropium bromide exhibits a pH-rate minimum at pH 3.5. Acid equivalent is given in units of normality which defines a pH range equivalent to 2.0-4.7 in an aqueous system (see pages 10-12). Table 2 (page 16) shows a formulation comprising ipratropium bromide, ethanol, water and ascorbic acid. Ascorbic acid is added in an amount of from 0.00015 to 5.0 mg/mI and the optimum pH level of the formulation is maintained at a range of from 2.0 to 4.7.

Bozung et al teach medicament compositions based on <u>anticholinergic</u> compounds which have a long-lasting effect and betamimetics, which have a long-lasting effect, processes for their production and their use in the therapy of respiratory ailments, especially **COPD** (col. 1, lines 11-16). **Tiotropium bromide monohydrate** is the preferred anticholinergic (col. 5, lines 51-55). The medicaments for inhalation are

dissolved in an aqueous or ethanolic solution, wherein solvent mixtures of ethanol

Jager does not specifically teach adding edetic acid or a salt thereof.

and water are also suitable. Other adjuvants, such as preservatives, e.g. benzalkonium chloride in concentration range of 8 to 12 mg/100 ml are added. Complex formers like **EDTA**, citric acid, ascorbic acid can be added. The medicament is present in an amount of from **0.001 to 5%** (see col. 6, line 39 to col. 7, lines 17-40).

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It would have been obvious to one of ordinary skill in the art at the time the invention was made to have modified the formulations comprising anticholinergics such as tiotropium and ipratropium, solvent, and acid with a pH level of from 2.0 to 4.7 as taught by Jager by implementing the acid of Bozung et al to prepare a similar formulation with acceptable and suitable stability. In other words, all the claimed elements were known in the prior art and one skilled in the art could have combined the elements as claimed by known methods with no change in their respective functions, and the combination would have yielded predictable results to one of ordinary skill in the art at the time of the invention.

Claims 38-49, 51, 52, 81-92, 94 and 95 are rejected under 35 U.S.C. 103(a) as being unpatentable over Freund et al or alternatively over Jager et al in view of Bozung et al as applied to claims listed above, and further in view of Weston et al (WO 9114468).

Freund et al, discussed above, lacks specific teachings on the inhalation device.

Jager et al in view of Bozung et al, discussed above lack specific disclosure of the inhalation device.

Weston et al discloses a metered dose inhaler which incorporates metering means for metering a quantity of fluid, and the atomizing means is provided by a mechanical break up device through which the metered quantity of fluid is passed to atomise it when it is subject to said increase in pressure (page 7, lines 5-9). For dispensing a spray of an aqueous solution of a medicament for inhalation into lungs, the droplet size is desirably less than 10 micrometers, typically 2 to 6 micrometers.

Weston also discloses that very high pressures can be generated in the pump cylinder or pressure and nozzle orifice diameters can be used, for example up to 100 micrometers, typically greater than 30 to 50 micrometers. The preferred pressures are from 50 to 400 bar, and more preferably from 100 to 350 bar with nozzle orifice of from 1 to 12 micrometers (page 12, lines 1-32).

It would have been obvious to a person of ordinary skill in the art at the time the invention was made to have utilized the preparation of Freund et al. or Jager et al and Bozung et al, by incorporating it in a device suitable for such preparations and because it is made simpler in design and cheaper to produce and suited to its function, as taught by Weston et al.

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## **Double Patenting**

The nonstatutory double patenting rejection is based on a judicially created doctrine grounded in public policy (a policy reflected in the statute) so as to prevent the unjustified or improper timewise extension of the "right to exclude" granted by a patent and to prevent possible harassment by multiple assignees. See *In re Goodman*, 11 F.3d 1046, 29 USPQ2d 2010 (Fed. Cir. 1993); *In re Longi*, 759 F.2d 887, 225 USPQ 645 (Fed. Cir. 1985); *In re Van Ornum*, 686 F.2d 937, 214 USPQ 761 (CCPA 1982); *In re Vogel*, 422 F.2d 438, 164 USPQ 619 (CCPA 1970); and, *In re Thorington*, 418 F.2d 528, 163 USPQ 644 (CCPA 1969).

A timely filed terminal disclaimer in compliance with 37 CFR 1.321(c) or 1.321(d) may be used to overcome an actual or provisional rejection based on a nonstatutory double patenting ground provided the conflicting application or patent either is shown to be commonly owned with this application, or claims an invention made as a result of activities undertaken within the scope of a joint research agreement.

Effective January 1, 1994, a registered attorney or agent of record may sign a terminal disclaimer. A <u>terminal disclaimer</u> signed by the assignee must fully comply with 37 CFR 3.73(b).

Claims 1-14, 16, 18-20, 22-31, 38-66, 68 and 70-95 are provisionally rejected on the ground of nonstatutory obviousness-type double patenting as being unpatentable over claims of copending Application No. 11/068,134 (US 20050147564). Although the conflicting claims are not identical, they are not patentably distinct from each other because the instant claims would have been anticipated by the reference claims. The claims of the co-pending application are drawn to a formulation comprising a first active agent comprising a tiotropium salt in a concentration range of between 0.0005% and 5% by weight, a steroid, a solvent such as water or ethanol and a preservative, wherein

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the formulation has a pH of from 2.0 to 3.5. The claims of instant application are drawn to a similar preparation. The difference is that the steroid is not required.

This is a <u>provisional</u> obviousness-type double patenting rejection.

Claims 1-14, 16, 18-20, 22-31, 38-66, 68 and 70-95 are provisionally rejected on the ground of nonstatutory obviousness-type double patenting as being unpatentable over claims of copending Application No. 10/392,558 (US 20040019073). Although the conflicting claims are not identical, they are not patentably distinct from each other because the instant claims would have been anticipated by the reference claims. The claims of the co-pending application are drawn to a formulation comprising a tiotropium salt in a concentration range of between 0.01 and 0.06 g per 100 ml of formulation, a solvent such as water and a preservative, wherein the formulation has a pH of from 2.7 to 3.1. The claims of instant application are drawn to a similar preparation. The difference is that the concentration range of tiotropium is slightly different.

This is a <u>provisional</u> obviousness-type double patenting rejection.

Claims 1-14, 16, 18-20, 22-31, 38-66, 68 and 70-95 are provisionally rejected on the ground of nonstatutory obviousness-type double patenting as being unpatentable over claims of copending Application No. 11/267,354 (US 20060057074). Although the conflicting claims are not identical, they are not patentably distinct from each other because the instant claims would have been anticipated by the reference claims. The claims of the co-pending application are drawn to a formulation comprising a first active

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agent comprising a tiotropium salt, a steroid, a betamimetic and a solvent such as water or ethanol. The preparation has a pH of from 2.0 to 7.0 (claim 17). The claims of instant application are drawn to a similar preparation. The difference is that the steroid and the betamimetic are not required.

This is a <u>provisional</u> obviousness-type double patenting rejection.

Claims 1-14, 16, 18-20, 22-31, 38-66, 68 and 70-95 are provisionally rejected on the ground of nonstatutory obviousness-type double patenting as being unpatentable over claims of copending Application No. 11/006,940 (US 20050148562). Although the conflicting claims are not identical, they are not patentably distinct from each other because the instant claims would have been obvious over the reference claims. Instant claims are drawn to formulations comprising an anticholinergic, preferably tiotropium and a second active agent such as a steroid. Formulations can be in a solution form and thus require a solvent. The preferred pH rang is from 2 to 7 (see e.g. claims 114 and 229). The claims of instant application are drawn to a similar preparation. The difference is that the second active agent such as steroid is not required.

This is a <u>provisional</u> obviousness-type double patenting rejection.

## Response to Arguments

Applicant's arguments filed 04/21/08 have been fully considered but they are not persuasive.

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Applicant argues that "the subject matter as a whole was not obvious in view of Freund et al. at the time the invention was made". Applicant continues that "the reduction in spray anomalies with low levels of edetic acid or edetic acid salt was not taught by Freund et al". Applicant refers to Table 1 on page 3 of Freund et al which shows a concentration of 50 mg/100 ml or more of EDTA in an ipratropium bromide solution yielded "0" spray anomalies as compared to tests run at lower levels of EDTA having between 2-6 spray anomalies. This is not persuasive because 1) Freund teaches addition of EDTA in a concentration range of from 10 to 100/100ml, which reads on the claimed range. 2) Spray anomalies are not a restriction or limitation of the instant claims. Instant claims are drawn to a preparation (product) and are examined based on their components and not how the product functions. 3) Where the general conditions of a claim are disclosed in the prior art, it is not inventive to discover the optimum or workable ranges by routine experimentation. In re Aller, 220 F. 2d 454, 105 USPQ 233 (CCPA 1955).

Applicant, by referring to MPEP 2144.05 (III) regarding overlapping ranges, argues that "by showing the criticality and unexpected results of the claimed ranges" the obviousness can be overcome. This is correct, however not persuasive here because a criticality has not been shown. Table 1 of Freund et al reference, shows 8 tests performed on formulations containing 0, 0.1, 1, 50 and 75 mg/100ml EDTA. While formulations containing from 0 to 1 mg/100ml EDTA showed spray anomalies, the formulations comprising 50 or 75 mg/100ml EDTA did not show any spray anomalies. This test is not commensurate with Applicant's arguments because there is no tests

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performed on 10mg or 25mg which would be more comparable with 50mg and 0-25mg/100ml range that is claimed here. Thus Table 1 of Freund et al is not a proper comparison for the claimed range. Applicant is also referring to the data provided with their arguments here (an attachment filed with their arguments of 04/21/08) and believes that the said data shows criticality. This is not persuasive because the data does not show criticality or unexpected results. At pH levels of from 2.7 to 3.0, the number of sprays at 0-50mg/100g NaEDTA are not consistent with any conclusion. For example, at pH of 2.7 and 2.8, the formulations comprising 10 mg of NaEDTA showed 0 number of sprays with deviation, but at pH of 3.0, the formulations comprising 25mg showed 0 number of sprays with deviation. The results for the pH level of 2.7 (at 10 and 25mg) were very similar to those with a pH of 3.2, which is outside of Applicant's optimum pH level. On the other hand the number of sprays with deviation for the formulations at pH level of 2.8 and 25mg were the same as those for the pH of 3.1 and 50 mg. Thus Applicants assertion that "An improvement of spray quality at lower pH values (2.7-3.0) in combination with lower NaEDTA concentrations (10 and 25 mg) is observed" is not found persuasive.

Applicant argues that "If a provisional nonstatutory obviousness-type double patenting rejection is the only rejection remaining in the earlier filed of the two pending applications,....the Examiner should withdraw that rejection...". However, the claims are not in condition for allowance at this time, thus the said Double Patenting rejections remain pending.

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Applicant's amendment necessitated the new ground(s) of rejection presented in this Office action. Accordingly, **THIS ACTION IS MADE FINAL**. See MPEP § 706.07(a). Applicant is reminded of the extension of time policy as set forth in 37 CFR 1.136(a).

A shortened statutory period for reply to this final action is set to expire THREE MONTHS from the mailing date of this action. In the event a first reply is filed within TWO MONTHS of the mailing date of this final action and the advisory action is not mailed until after the end of the THREE-MONTH shortened statutory period, then the shortened statutory period will expire on the date the advisory action is mailed, and any extension fee pursuant to 37 CFR 1.136(a) will be calculated from the mailing date of the advisory action. In no event, however, will the statutory period for reply expire later than SIX MONTHS from the date of this final action.

Any inquiry concerning this communication or earlier communications from the examiner should be directed to MINA HAGHIGHATIAN whose telephone number is (571)272-0615. The examiner can normally be reached on core office hours.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, Johann Richter can be reached on 571-272-0646. The fax phone number for the organization where this application or proceeding is assigned is 571-273-8300.

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/Mina Haghighatian/

Mina Haghighatian Primary Examiner Art Unit 1616